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- (71) Applicant (for all designated States except US):
  PHARMAPRODUCTS UK LIMITED [GB/GB];
  7th Floor, Castle Chambers, 43 Castle Street, Liverpool,
  Merseyside L2 9TL (GB).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): SALVAGGIO, Antonio [IT/IT]; Via Cesare Battisti, 35, I-95021 Acicastello (CT) (IT). NICOLETTI, Pierferdinando [IT/IT]; Via Luigi Sturzo, 3, I-95021 Cannizzaro (CT) (IT). MACRI', Battesimo [IT/IT]; Polo Universitario dell'Annunziata, c. da SS Annunziata, I-98168 Messina (IT).

- (74) Agents: MINOJA, Fabrizio et al.; Bianchetti Bracco Minoja S.r.l., Via Rossini, 8, I-20122 Milano (IT).
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WO 03/101477 PCT/EP03/05551

## USE OF UK114 IN THE TREATMENT OF LEISHMANIASIS

This invention relates to the use of the protein UK114, possibly associated with ubiquitin, to treat leishmaniasis in humans and animals.

The protozoa of the *Leishmania* genus are intracellular parasites of the macrophages and dendritic cells of the dog, man and numerous wild animals. On the basis of the classification criteria used in human medicine, leishmaniasis presents in three clinical forms: visceral (known in man as "kala-azar"), cutaneous and mucocutaneous.

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In the case of the dog, a cutaneous and a visceral form were separately classified in the past because of the characteristic clinical picture, but they are now both regarded as progressive forms of the same disease, known as "generalised canine leishmaniasis".

The vector is a sand-fly of the *Phlebotomus* genus in the Old World and the *Lutzomyia* genus in the New World. The protozoa multiply in the sand-fly and are transformed into infectious organisms.

Parasites of the Leishmania genus appear as rounded or oval organisms in the macrophage, with the rod-like kinetoplast adjacent to the nucleus. The organism, which measures 2 to 5  $\mu$ m in diameter, possesses a rudimentary flagellum that does not extend beyond the edge of the cell. This amastigote form of the parasite is ingested by the sand-fly during the blood meal. The protozoon is transformed in the intestine of the intermediate host into the promastigote form, characterised by a long free flagellum that protrudes from the anterior extremity of the parasite. The organism has an elongated shape and can grow to a length of 15  $\mu$ m, excluding the flagellum, which usually has the same dimensions as the body.

The amastigotes ingested reach the intestine of the sand-fly, where they are transformed into promastigotes. The promastigotes divide repeatedly by

WO 03/101477 PCT/EP03/05551

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binary fission, and subsequently migrate in the anterior direction. In the pharynx, the parasites turn into highly mobile metacyclic promastigotes, which migrate towards the proboscis. The promastigotes are transmitted to the new vertebrate host by means of the sand-fly's bite.

In the vertebrate host, the promastigotes are ingested by the monocytes/macrophages. After being ingested, the promastigote turns into an amastigote. The amastigotes divide by binary fission in the parasitophorous vacuole until their number is sufficient to rupture the macrophage. The amastigotes thus released are ingested by other macrophages.

The ability of the amastigotes to survive in the macrophages and spread throughout the body depends on factors intrinsic to the parasite and on factors associated with the type of cell-mediated immune response developed by the host. If parasitic macrophages are sufficiently stimulated by T-helper (Th) lymphocytes, they produce numerous lysosome enzymes and other factors including oxygen metabolites, hydrogen superoxide and peroxide and nitrous oxide (NO), which are toxic to the parasite.

The type of cell-mediated immune response and interleukin (IL) profile produced determine resistance or sensitivity to *Leishmania* infection. In laboratory animals, resistance to *Leishmania* infection is characterised by the **Th1** response, with production of IL-12 and interferon gamma (IFNγ) and activation of the macrophages which eliminate the parasite. Conversely, in animals sensitive to infection, the response is type **Th2**, characterised by production of IL-4 and IL-10 with consequent suppression of the parasiticidal activity of the macrophages and stimulation of the B lymphocytes with an increase in the production of immunoglobulins.

The humoral immune response in leishmaniasis is impressive, but not protective. The specific antibodies produced against *Leishmania* have no neutralising action against the parasite.

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characterised by:

In animals sensitive to the disease the protozoon spreads throughout the body, in the macrophages. The parasite has been observed in all the organs and tissues of the body except the central nervous system. Slow, continuous contact between the parasitic antigen and the immunocompetent cells forms the basis for the pathogenetic development of the disease, which is

hyperglobulinaemia, generally polyclonal, associated with continual stimulation of the B lymphocytes, which causes an increase in total proteins and inversion of the albumin/globulin ratio;

production of auto-antibodies, probably due to a cross-reaction between parasitic antigens and self-antigens, causing thrombocytopenia and anaemia;

production and deposit of immunocomplexes responsible for the vasculitis, glomerulonephritis and polyarthritis syndromes.

The pathogenesis of the skin lesions present in most sufferers is not yet clear. According to some authors, the persistence of the parasite in the macrophage continually stimulates infiltration by inflammatory cells, especially plasma cells, macrophages and lymphocytes, into the dermis. According to other authors, the deposit of immunocomplexes is the main cause of dermatitis which, on histological examination, often presents lesions similar to those caused by other diseases induced by immunocomplexes, such as systemic Lupus erythematosus. Finally, the skin alterations may be the result of vasculitis.

The symptoms of canine leishmaniasis are highly variable, and may include peripheral lymphadenopathy (over 90% of infected subjects), skin lesions (>80%), chronic conjunctivitis (50%), onychogryphosis (40%), anorexia (>35%), increased appetite (30%), weight loss (30%), fever (20%), kidney failure (20%), epistaxis (10%), uveitis (8%) and gait disorders (6%).

The skin signs are among the most important in the disease. Various

WO 03/101477 PCT/EP03/05551

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types of macroscopic and microscopic lesions have been described in canine leishmaniasis: dry exfoliative dermatitis, ulcerative dermatitis, nodular dermatitis, sterile pustular dermatitis, paronychia, and nasal and/or digital hyperkeratosis. The skin lesions are generally chronic, symmetrical and not itchy.

Recent studies clearly demonstrate the existence of a TNF-independent compensation mechanism able to activate the macrophages in the anti-leishmania response. As the **Th1** response mediated by interleukin-12 (IL-12) and interferon gamma (IFN $\gamma$ ) is paradoxically responsible not only for activation of the macrophages, but also for nearly all the symptoms of leishmaniasis, it may be advantageous to boost this compensation mechanism by inhibiting the **Th1** response.

The current elective treatment, based on antimony gluconate administered by infiltration (in cutaneous l.) or injection (in the other forms) can cause toxic effects (nausea and vomiting) sufficiently serious to require discontinuance of the treatment, which is replaced by treatment with aromatic diamines such as pentamidine, whose tolerability is generally poor.

It has now been found that the protein with molecular weight 14 kDa in SDS-PAGE, obtainable by extraction from mammal liver with perchloric acid, called UK114 and disclosed in EP 574394 and US 5792744, is useful in the treatment of leishmaniasis, possibly associated with ubiquitin (UK110).

Recombinant protein UK114 is known from WO 00/63368.

Subcutaneous administration of UK114 and ubiquitin to a group of 10 dogs with manifest clinical symptoms of leishmaniasis (peripheral lymphadenopathy and skin lesions of a high degree, mainly represented by sores and bleeding ulcers with loss of substance, anorexia and weight loss), at the doses and times indicated in the table, led to complete healing of all the lesions during the treatment period.

No adverse effects were observed during the treatment.

These findings demonstrate that the administration of UK114 and ubiquitin cures the clinical symptoms of leishmaniasis in a totally safe manner. This is very interesting in view of the high toxicity of the drugs currently used in treatment, and the possibility of a response that is sometimes unsatisfactory from the clinical standpoint.

**TABLE** 

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Patient's weight < 10 kg	1mg/day subcutaneously for 6 days 7th day: rest 1 mg/day subcutaneously for 6 more days
Patient's weight > 10 kg	As above, but doubling the dose: 2 mg/day

According to the invention, protein UK114 of extractive or recombinant origin will therefore be opportunely administered to subjects suffering from leishmaniasis by the parenteral route, in particular subcutaneously or intramuscularly, at doses ranging between approx. 0.5 and 10 mg a day, until the disappearance or substantial reduction of the symptoms. The compositions according to the invention, in the form of solutions or suspensions in preferably aqueous sterile solvents, may also contain ubiquitin in a quantity corresponding to 0.1-5 mg per unit dose.

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### **BIBLIOGRAPHY**

- 1. Urquhart G.M., Armour J., Duncan J.L., Dunn A.M, Jennings F.W. Veterinary Parasitology 2<sup>nd</sup> ed., Italian edition edited by C. Genchi, UTET 1998.
  - 2. Murray H.W., Jungbluth A., Ritter E., Montelibano C., Marino M.W. Visceral Leishmaniasis in Mice Devoid of Tumor Necrosis Factor and Response to Treatment, Infection and Immunity, November 2000, p. 6289-6293, Vol. 68, No. 11.
- 10 3. Katzung B.G. Basic & Clinical Pharmacology, 1995 by Appleton & Lange, E.N., Connecticut.

## **CLAIMS**

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- 1. Use of the protein UK114, possibly in combination with ubiquitin, for the preparation of pharmaceutical or veterinary compositions for the treatment of leishmaniasis in humans and animals.
- 2. Pharmaceutical or veterinary compositions containing the extractive or recombinant protein UK114, possibly associated with ubiquitin, together with a suitable vehicle, for the treatment of leishmaniasis in humans and animals.

### **INTERNATIONAL SEARCH REPORT**

Interioral Application No
PCT/EP 03/05551

A. CLASSIF IPC 7	FICATION OF SUBJECT MATTER A61K38/17 A61P33/02									
According to	According to International Patent Classification (IPC) or to both national classification and IPC									
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	ion searched other than minimum documentation to the extent that su									
	ata base consulted during the International search (name of data bas ternal, WPI Data, PAJ, BIOSIS, EMBAS									
C. DOCUME	ENTS CONSIDERED TO BE RELEVANT									
Category °	Citation of document, with Indication, where appropriate, of the rele	evant passages	Relevant to claim No.							
Х	WO 98 42366 A (MERONI PIER LUIGI SPA (IT); PANERAL ALBERTO (IT); B 1 October 1998 (1998-10-01)	2								
A	page 1, line 20 - line 24 page 2, line 11 - line 16 page 4, line 1 - line 5		1							
		-/	·							
X Furti	her documents are listed in the continuation of box C.	Patent family members are listed	in annex.							
"A" docume consider filling of the consider of the country of the	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another or or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but than the priority date claimed	T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention.  X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone.  Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.  &' document member of the same patent family								
	actual completion of the international search	Date of mailing of the international se	earch report							
	September 2003 mailing address of the ISA	26/09/2003 Authorized officer								
· ·anie and	Funding address of the 13-december 2  European Patent Office, P.B. 5818 Patentiaan 2  NL – 2280 HV Rijswijk  Tel. (+31-70) 340–2040, Tx. 31 651 epo ni,  Ear. (+31-70) 340–3016	Hars, J								

## INTERNATIONAL SEARCH REPORT

int Pilonal Application No PCT/EP 03/05551

		101/21 03/03331
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	Relevant to claim No.
Category °	Citation of document, with Indication, where appropriate, of the relevant passages	
Х	PANERAL A E ET AL: "CHRONIC ADMINISTRATION OF UK-114, A MULTIFUNCTIONAL EMERGING PROTEIN, MODULATES THE TH1/TH2 CYTOKINE PATTERN AND EXPERIMENTAL AUTOIMMUNE DISEASES" ANNALS OF THE NEW YORK ACADEMY OF SCIENCES, NEW YORK ACADEMY OF SCIENCES, NEW YORK, NY, US, vol. 876, 1999, pages 229-235, XP000971426 ISSN: 0077-8923	2
Α	10011. 007. 0520	1
-	abstract page 230, paragraph 1 - paragraph 2 page 233, paragraph 1 page 234, last paragraph	
X	NICOLETTI FERDINANDO ET AL: "Prevention and treatment of lethal murine endotoxemia by the novel immunomodulatory agent MFP-14."  ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, vol. 45, no. 5, May 2001 (2001-05), pages 1591-1594, XP002253826	
	ISSN: 0066-4804	1
A	abstract page 1591, left-hand column -right-hand column, paragraph 2	
X	WO 99 43340 A (ZETESIS SPA ;SANTI CESARE (IT); BARTORELLI ALBERTO (IT)) 2 September 1999 (1999-09-02) page 1 claim 4	2
х	WO 00 78329 A (ZETESIS SPA ;PANERAI ALBERTO (IT); BARTORELLI ALBERTO (IT); NICOLE) 28 December 2000 (2000-12-28) claim 1	2
Α	DATABASE BIOSIS 'Online! BIOSCIENCES INFORMATION SERVICE, PHILADELPHIA, PA, US; July 2001 (2001-07) KHASKHELY NOOR MOHAMMAD ET AL: "Pre-exposure with low-dose UVA suppresses lesion development and enhances Th1 response in BALB/c mice infected with Leishmania (Leishmania) amazonensis." Database accession no. PREV200100345475 XP002253827 abstract & JOURNAL OF DERMATOLOGICAL SCIENCE, vol. 26, no. 3, July 2001 (2001-07), pages 217-232, ISSN: 0923-1811	1,2
	133N. V323 1011	

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
WO 9842366	A	01-10-1998	IT	MI970694 A1	25-09-1998
WU 3042300	^	01 10 1990	ĀŪ	7207298 A	20-10-1998
			BR	9808625 A	16-05-2000
			CN	1251043 T	19-04-2000
			MO	9842366 A1	01-10-1998
			EP	0969859 A1	12-01-2000
			ΗÜ	0002304 A2	28-11-2000
•			JP	2001518107 T	09-10-2001
			NO	994622 A	23-09-1999
			NZ	337985 A	23-02-2001
			PL	335829 A1	22-05-2000
			TR	9902331 T2	21-01-2000
			ÜS	6255283 B1	03-07-2001
WO 9943340 A	Α	02-09-1999	IT	MI980356 A1	24-08-1999
			ΑT	208627 T	15-11-2001
			AU	2624499 A	15-09-1999
			CA	2321717 A1	02-09-1999
			DE	69900466 D1	20-12-2001
			DE	69900466 T2	04-04-2002
			DK	1061938 T3	11-02-2002
			WO	9943340 A1	02-09-1999
			EP	1061938 A1	27-12-2000
			ES	2165727 T3	16-03-2002
			JP	2002504517 T	12-02-2002
			PT	1061938 T	29-04-2002
WO 0078329	Α	28-12-2000	IT	MI991384 A1	22-12-2000
			ΑU	5221700 A	09-01-2001
			BR	0011832 A	05-03-2002
			CA	2377639 A1	28-12-2000
			CN	1399555 T	26-02-2003
			CZ	20014631 A3	17-04-2002 28-12-2000
			MO	0078329 A2	10-04-2002
			EP	1194158 A2	28-09-2002
			HU	0201796 A2	28-09-2002
			JP	2003502380 T	21-12-2001
			NO	20016326 A	25-07-2003
			NZ	516159 A 200103748 T2	21-05-2002
			TR ZA	200103748 12 200110213 A	12-12-2002
			44	COULTOTIO W	12 12 2002